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(54) Title: Continuous preparation of isocyanates

(57) Abstract: The invention relates to a process for preparing polyisocyanates by reacting organic amines with phosgene, wherein the reaction is carried out in at least three stages, with the first stage being carried out in a mixing apparatus, the second stage in at least one residence apparatus and the third stage in at least one separation apparatus and the pressure in each successive stage being lower than that in the previous stage.

Description

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relates to process for present invention a or aliphatic isocyanates. preparing aromatic is preference given aromatic isocyanates, (MDI) and methylenedi (phenyl isocyanate) in the case of aliphatic (TDI), while diisocyanate given hexamethylene preference is to isocyanates, diisocyanate (HDI) and isophorone diisocyanate (IPDI) and others.

It is an object of the present invention to develop a two-stage or multistage process which gives isocyanates in very high chemical yields and with high space-time yields at a low phosgene holdup.

In the process of the present invention, the reaction between organic amine and phosgene is carried out in two or more stages in an inert solvent, preferably toluene, chlorobenzene, dichlorobenzene or a mixture of 20 latter two, and using an excess of phosgene, wherein the pressure is reduced between each of the stages and the first phosgenation stage comprises a static mixer, preferably a nozzle, the second stage comprises a residence apparatus and the third stage 25 comprises a (reaction) column. The pressure upstream of the nozzle is preferably 3-70 bar, more preferably 15-45 bar. The residence reactor of the second stage is pressure of 2.5-35 bar, preferably operated at a preferably 15-35 bar. Downstream of the nozzle, 30 pressure is reduced to the pressure of the second stage by means of a valve or some other device suitable for this purpose. It is, however, also possible to use the natural pressure drop in the nozzle for reducing the pressure. The reactor of the first stage, usually a 35 static mixer, can also be integrated into the reactor of the second stage, viz. a residence apparatus. The

reactor of the third phosgenation stage, usually a column, preferably a reaction column, is preferably operated at a pressure of 2-20 bar, preferably 3.5-16 bar. Downstream of the reactor of the second stage, the pressure is reduced to the pressure of the reactor of the third stage by means of a valve or some other device suitable for this purpose. Once again, a natural to achieve may be sufficient drop pressure reduction in pressure. As reactor of the third stage, preference is given to using a (reaction) column as is described, for example, in WO 99/54289 (DE 19817691).

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The preparation of isocyanates from amines and phosgene is known. The reaction is, depending on the type of amine, carried out either in the gas phase or in the 15 liquid phase, either batchwise or continuously Siefken. Liebigs Ann. 562, 75 (1949)). The continuous preparation of organic isocyanates by reaction of primary organic amines with phosgene has been described many times and is carried out on a large industrial 20 scale (cf., for example, Ullmanns Enzyklopädie der Technischen Chemie, Volume 7 (Polyurethane), 3rd revised edition, Carl Hanser Verlag, Munich-Vienna, p. isocyanates TDI (tolylene (1993).). The aromatic diisocyanate) and MDI (methylenedi(phenyl isocyanate)) 25 (polymethylene-polyphenylene polyisocyanate) latter two and the aliphatic and mixtures of the (hexamethylene diisocyanate) isocyanates HDI isophorone diisocyanate (IPDI) particular in are prepared industrially around the world. 30

industrial syntheses of the aromatic Present-day and the aliphatic TDI diisocyanates MDI and diisocyanates HDI and IPDI are carried out virtually exclusively in continuous processes. A continuous process for carrying out the reaction in a plurality of vessels through which the reaction mixture

continuously is described, for example, in DE 844896. In general, the continuous embodiment of the process is carried out in two stages. In the first stage of the phosgenation, the amine is reacted with phosgene to form the corresponding carbamoyl chloride and hydrogen chloride and the amine hydrochloride. The is very fast, strongly between amine and phosgene exothermic and proceeds even at very low temperatures. To minimize by-product and solids formation, amine and phosgene, each in an organic solvent if appropriate, 10 therefore have to be mixed quickly, which is why the first phosgenation stage is generally carried out in a mixing apparatus, preferably a nozzle. The second stage of phosgenation comprises both the decomposition of the carbamoyl chloride to form the desired isocyanate and 15 hydrogen chloride and the phosgenation of the amine form the carbamoyl chloride. The hydrochloride to phosgenation temperature of the second stage is generally higher than that of the first.

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The reaction of amine and phosgene in the liquid phase is very fast at all industrially customary temperatures and pressures. For this reason, good mixing of the reactants is sought, in order to suppress secondary reactions. The phosgenation of primary amines in a mixing reactor as first stage of the phosgenation has been described many times.

Mixing apparatuses can basically be classified as dynamic mixers, e.g. stirrers and turbines, and static mixers, rotor-stator systems such as Kenics mixers, Schaschlik mixers, SMV mixers, and jet mixers such as nozzles or T-mixers (Chem. Ing. Tech. MS 1708/88, Fortschr. Verf. Technik 23, 1985, 373, Ind. Eng. Chem. Res. 26, 1987, 1184).

Known mixing apparatuses include, in particular,

nozzles such as annular slit nozzles (DE 1792660), nozzles (DE 3744001), smooth-jet mixing annular hole nozzles (EP 0065727), fan jet nozzles (DE 2950216), angled jet chamber nozzles (DD 300.168), three-stream nozzles (DD 132340), countercurrent mixing chambers nozzles (FR 69428), fluid backup (DE-C 1146872), Venturi mixing nozzles (DE-B 1175666). In-line mixers (US 3321283), centrifugal mixing pumps or reaction (EP 0291819), tubular reactors mixing pumps 3226410) or microstructure mixers (EP 0928785) are also known. CA 832432 describes the use of ultrasound waves for mixing.

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EP 0830894 describes a mixing reactor for the phosgenation of primary amines, in which the inlet for one substance is located in the axis of the mixing chamber and the inlet for the (at least one) other substance is configured as a multiplicity of nozzles arranged rotationally symmetrically around the axis of the mixing chamber, with each of these nozzles having a pin which is movable in the direction of the nozzle axis and can free the nozzle of adhering solids.

DD 132340 describes a process for the phosgenation of form monoisocyanates, diisocyanates amines polyisocyanates under superatmospheric pressure and at elevated temperature in the presence of a uniform solvent, in which an amine/monochlorobenzene mixture and a phosgene/monochlorobenzene mixture are parallel as a plurality of substreams into a reactor, with part of the phosgene/monochlorobenzene centrally and introduced amine/monochlorobenzene mixture being introduced around this central stream and being in turn enclosed by a polyamine/ phosgene/monochlorobenzene mixture. The monochlorobenzene mixture is, for example, fed as an annular stream into the phosgenation reactor at 150°C.

Before entry into the reactor, a rotating motion is imparted to the mixture by means of an appropriate device. In and around the twist-inducing mixture, polyamine/monochlorobenzene а phosgene/monochlorobenzene mixture which has been heated to 150°C is fed as reactant into the reactor. The relative velocity of the two reactants is about 15 m/s.

phosgenation stage, which may the second simultaneously be used as phase separation vessel, a 10 apparatuses has also of multiplicity established. The preparation of isocyanates from the corresponding amines by phosgenation is carried out in stirred vessels (e.g. DE-A 1468445), in cascades stirred vessels (DE-C 844896), in packed reaction 15 columns (e.g. WO 99/54289) or in unpacked columns (e.g. Encyclopädie der technischen Chemie, Ullmanns edition (1977), pages 351 ff). A circulating mode of operation is also realized by use of loop reactors in order to provide a sufficient residence time for 20 complete conversion at a limited reaction volume (and holdup).

first stage of the isocyanate synthesis is The frequently carried out at a very low temperature and 25 the second stage at significantly higher temperature in residence apparatus. This mode of operation frequently described as cold-hot phosgenation and is described, for example, in W. Siefken, Liebigs Analen der Chemie 562 (1949), page 96. Firstly, a suspension 30 the intermediates carbamoyl chloride and amine temperature, low is prepared at hydrochloride particular at 0°C or room temperature, but at most 60°C, and this is then converted into the isocyanate at higher temperatures, in particular in the range 100-35 Such apparatus. two-stage 200°C, in a residence in the Kunststoffhandbuch, described processes are

Volume 7 (Polyurethane), 3rd revised edition, Carl Hanser Verlag, Munich-Vienna, p. 76ff (1993), and, for example, in the patents DE 2058032, DE 2153268, DE 2908703, DE 1233854.

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DE 949227 describes a cold-hot phosgenation process for the continuous preparation of isocyanates by reaction of amines with phosgene in the liquid phase in the which, solvent, in in the of а presence phosgenation, a solution or slurry of the amine in an inert solvent is continuously combined with phosgene or a solution of phosgene in an inert solvent a mixing apparatus with intensive stirring and without external cooling in the cold phosgenation and the reaction mixture obtained in this way is then phosgenation. As subjected to the hot and centrifugal turbomixers apparatuses, pumps mixing apparatuses having moving mechanical parts in general are claimed. The residence time in the mixing apparatus ranges from a few seconds to one minute.

DE 949228 describes a cold-hot phosgenation process for continuous preparation of monocyclic aromatic diisocyanates (e.g. tolylene diisocyanate), in which a suspension is prepared continuously from the parent amine of the isocyanate and the total amount of the solvent used in the phosgenation, the suspension is continuously reacted cold with phosgene, the reaction product is subsequently pushed continuously through one or more vertical or slanted tubes known as phosgenation if desired with towers in which the material is, introduction of qaseous phosgene, heated the solution the and phosgenation temperature subsequently freed of dissolved phosgene by stripping а column. stream in with а dry inert qas solvent. The cold Dichlorobenzene as was used phosgenation is carried out at 0°C in a stirred vessel,

the reaction mixture is then preheated to 30°C and finally reacted at 170°C in the hot phosgenation in two phosgenation towers connected in series to give the introduced isocyanate. Gaseous phosgene is bottom of the second phosgenation tower. The hydrogen 5 chloride/phosgene/solvent mixture taken off at the top of the two phosgenation towers is partially condensed respective condenser the the at in respective phosgenation tower and is recirculated to the bottom. The incondensible gas comprising phosgene 10 and hydrogen chloride is passed to a phosgene/hydrogen chloride separation unit or to disposal. The liquid and completely phosgenated reaction product leaving the second phosgenation tower runs into a bubble cap tray column and is there freed of dissolved phosgene by 15 stripping in countercurrent with a stream of nitrogen. The vapor taken off at the top is passed to a condenser and condensed solvent is recirculated to the top of the At the bottom of the column, the reaction product in the solvent is taken off and passed to 20 distillation.

DE 952086, too, describes a cold-hot phosgenation. The hot phosgenation is carried out in upright reaction towers filled with Raschig rings or other packing elements. The cold phosgenation is carried out at 0°C and the hot phosgenation is carried out using a rising temperature profile from 120°C to 160°C.

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In DE 958558, the circulated solvent is introduced not at the lower end of the reaction towers but in the cold phosgenation stage. Dilution of the reaction product from the cold phosgenation gives the advantage of a solution having a relatively low viscosity instead of a viscous suspension of carbamoyl chloride and amine hydrochloride.

DE 2058032 also describes a cold-hot phosgenation. The hot phosgenation is carried out in horizontal tube reactors at up to about 200°C, and the reaction mixture from the cold phosgenation is fully reacted in the hot phosgenation stage with continual mechanical mixing and a slowly rising temperature profile. The cold phosgenation is carried out at 0°C.

US 2908703 describes a two-stage process for preparing aromatic isocyanates, in which the first reaction step 10 is carried out at 60-90°C, preferably 70-85°C, using chlorobenzene as solvent, and the second reaction step is carried out at a temperature which is high enough to decompose the intermediate to form the isocyanate. A solution of amine in an organic solvent, preferably 15 o-dichlorobenzene, and chlorobenzene orphosgene are introduced simultaneously into a stirred and heated reactor so that a saturated phosgene solution is formed and the phosgene excess is at least 50% over the stoichiometric amount of phosgene for the 20 reaction reaction. The mixture amine/phosgene subsequently heated to the decomposition temperature of the carbamoyl chloride and amine hydrochloride. finally separated off formed is isocyanate other methods. distillation orfractional 25 batch procedure, the process alternative to this described can also be carried out continuously by carrying out the first reaction step in a first reactor and the second step in a second reactor. The reaction is generally carried out at atmospheric pressure. The 30 concentration of the amine in the organic solvent is 2-20% by weight, preferably 5-10% by weight. concentrations lead to formation of by-products, particular ureas and polyureas.

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In US 3381025, the phosgenation of an organic primary amine to form the corresponding isocyanate is carried

out in two stages at <60°C in the first stage and 100-190°C in the second stage. A mixture of the inert solvent, excess phosgene and the hydrogen chloride formed is taken off from the second reaction stage and the hydrogen chloride is separated off from this mixture by cooling the mixture to -20°C. The cold liquid mixture of phosgene and solvent obtained is recirculated to the first reaction stage.

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- DE 2153268 describes a process for a continuous cold 10 phosgenation of organic primary amines by reaction with phosgene in an inert solvent in a solution of not self-priming centrifugal multistage, centrifugal pump at the same time conveys the resulting reaction mixture to the subsequent hot phosgenation 15 stage. The phosgene solution enters the pump at from -105°C to +25°C and the amine solution enters it at from 50 to 100°C. The reaction mixture leaves the pump at The concentration of the amine from 50 to 110°C. solution is from 5 to 40% by weight, and that of the 20 phosgene solution is 20-65% by weight. The amount of phosgene is at least 1 mol, preferably 1.5-3 mol, per mol of amine group.
- A disadvantage of the two-stage procedure with a low 25 temperature in the first stage and a high temperature in the second stage (cold-hot phosgenation) is the low reaction rates and thus low space-time yields owing to the low temperatures in the first phosgenation stage. The low temperatures (high solubility of phosgene) and 30 the long reaction times (large reactors) additionally imply a high phosgene holdup, which is undesirable from a safety point of view. Low temperatures are also problematical because of the massive precipitation of chloride which is formed as an carbamoyl 35 the decomposes rapidly at intermediate and temperatures. This brings with it the risk of blockages

caked material. Furthermore, and the formation of cooling of the reactants and later heating of the reaction mixture is energetically disadvantageous. achieve economical space-time yields, operation elevated temperature in all stages is necessary 5 industrial processes for preparing organic isocyanates by phosgenation of primary organic amines. However, the solubility of the phosgene in the reaction mixture and available for the phosgene excess therefore the reaction decreases at high temperatures, since the 10 reaction generally takes place in the liquid phase. A high excess of phosgene is nevertheless necessary to of isocyanate. yields high achieve describes the influence of pressure and temperature on the reaction and the phosgene excess. The lowering of 15 phosgene excess at elevated temperatures generally countered by means of an increased reaction pressure. DE-A 1768439 describes a process for the continuous preparation of organic isocyanates which employs a combination of high temperature above 180°C 20 and a high pressure of from 20 to 150 atm together with a high phosgene concentration in the reaction zone. The amount of amine introduced is from 2.5 to 5.5 times the stoichiometric amount. As a result of the extremely high pressure and the very high temperature, acceptable 25 space-time yields can still be achieved. The residence time of the reactants in the reaction zone is 5-60 s. The preferred solvent is chlorobenzene. A disadvantage of the process is the reduced yield and quality caused formation of by-products, the increased 30 particular ureas, as a result of the high temperature. In addition, industrial pressure apparatuses are very expensive and are problematical because of the high toxicity of phosgene. Carrying out the reaction of amine with phosqene at high pressure also has the 35 disadvantage that, in accordance with Henry's law, not only the phosgene concentration but also the hydrogen

concentration in the liquid phase is chloride formation of amine increased increased. The hydrochlorides is, however, undesirable since their according to generally accepted phosgenation is, opinion, very slow and therefore represents the ratedetermining step of the overall reaction. This leads to very long residence times and large phosgene holdups.

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EP 0065727 describes a process using a nozzle and tube reactor. A process for the continuous preparation of 10 monoisocyanates and polyisocyanates organic single-stage reaction by continuous combination solutions of primary monoamines or polyamines in inert amounts of phosqene excess organic solvents with dissolved in an inert organic solvent at pressures of 15 from 10 to 1000 bar, preferably from 25 to 150 bar, and temperatures of from 120 to 300°C, preferably from 150 250°C, in a mixing zone and, if appropriate, downstream reaction zone and continuous work-up, with the phosgene solution, which is used in excess, being 20 continuously placed in a mixing zone and the amine component, which is used in a deficiency, injected by means of a smooth-jet nozzle, is described. The smooth-jet nozzle which is essential to the process has an internal diameter of 0.1-30 mm. A differential 25 pressure of at least 0.5 bar, preferably 1-200 bar, in particular 3-50 bar, is maintained in the phosgene solution injected. The molar ratios of phosgene to amino groups are from 2:1 to 30:1, preferably from 3:1 to 18:1. The after-reaction zone can be a tube reactor, 30 a shell-and-tube reactor or, for example, a cascade of stirred vessels. The mean residence time in the mixing vessel and in the downstream reaction zone is from 5 seconds to 5 minutes. The reaction mixture leaving the after-reaction zone is depressurized to atmospheric 35 pressure in one or more stages in a depressurization vessel, resulting in a temperature drop of 50-150°C. Gas

and liquid phases are separated in the depressurization vessel. As solvents, preference is given to using chlorobenzene or o-dichlorobenzene.

GB 827376 describes a continuous process for preparing 5 aromatic isocyanates by reacting an amine in free form in a solvent or as a salt which readily decomposes to the amine and is suspended in this solvent with a solution of phosgene in an inert solvent at a pressure greater than $3 \cdot 10^5 \text{Pa}$, with the reactants being 10 introduced simultaneously with mixing into the lower end of a vertical tube reactor in which the reaction products rise rapidly to the upper end. The liquid phase is collected in a container from which it is taken off to isolate the isocyanate. This container can 15 be a phase separation apparatus which is operated under the same pressure, is connected via an overflow tube to the liquid outlet and has a throttle valve in the liquid outlet. The liquid collected in the container is fed into a column which is operated under atmospheric 20 at superatmospheric pressure and so that residual phosgene and hydrogen temperature, chloride are separated off at the top in gaseous form. The excess phosgene is condensed out (preferably by the hydrogen cooling water) from 25 means off mixture separated in chloride/phosqene container, and the hydrogen chloride which has been way is depressurized separated off in this discharged. The reactants are fed by means of one joint pump or two separate pumps into the tube reactor or 30 else mixed in a Venturi mixing nozzle, preferably one having separate inlets for the two reactants, and from there introduced into the tube reactor. The temperature in the tube reactor is given as 80-200°C, the pressure is greater than $3 \cdot 10^5$ Pa, at most the vapor pressure of 35 the reaction mixture and preferably $15-20\cdot10^5$ Pa.

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US 3226410 describes a continuous process for preparing aromatic isocyanates by mixing a stream of an aromatic amine into a phosgene stream in a tube reactor at Reynolds numbers of greater than 2100 (preferably 5000-2000000) and temperatures of 60-90°C, preferably 80least The amount of phosgene is at preferably from 6 to 12 mol, per mol of amine. The then, reaction solution is if appropriate preheating, fed into a second reactor, in particular a vessel or a column, which is at from 110 to 135°C, preferably from 110 to 120°C. The amine concentration is from 2 to 25% by weight, preferably from 5 to 10% by weight, and the phosgene concentration is from 10 to 100% by weight, preferably from 10 to 60% by weight. The pressure at which the phosgene stream is introduced into the tube reactor is 50-170 psig; the pressure of the amine stream has to be greater in order to prevent backmixing. The liquid phase comprising isocyanate, amounts of by-products, small solvent, relatively hydrogen chloride and phosgene dissolved in a solvent is taken off from the second reactor separately from the gas phase comprising hydrogen chloride, solvent, phosgene and traces of the isocyanate. Solvents used are chlorinated hydrocarbons which are inert and have a boiling point lower than that of the isocyanate. Particular preference is given to chlorobenzene.

The second reactor, which has a pressure of 45 psig or higher, is followed by a residence vessel and a buffer vessel from which the liquid phase is conveyed under level control to a column for removing excess phosgene. Phosgene, hydrogen chloride and solvent are taken off at the top and recirculated to the phosgene container. The bottom product comprising isocyanate and solvent is conveyed to a distillation, preferably a single-stage distillation, to separate off the solvent. The solvent which has been separated from the isocyanate is used

for absorption of the remaining phosgene from the hydrogen chloride stream. The phosgene taken off in the second reactor and in the buffer vessel is condensed in two stages and recirculated to the phosgene container. The uncondensed phosgene/hydrogen chloride mixture is conveyed to a phosgene absorber into which solvent recovered in the solvent separation is fed. The gas which has not been absorbed, mainly hydrogen chloride, is subsequently reacted with water in an absorber to form aqueous hydrochloric acid.

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The tube reactor should be constructed as a plug flow reactor without deflection plates, pockets or other internals which can produce dead zones so that settling of solids is prevented. The high Reynolds numbers and the design of the reactor as straight tubes are intended to lead to the liquid continually keeping the walls free of caked material.

DE 952086 describes a process for preparing isocyanates 20 from primary amines or their salts and phosgene, which the reactants are, in the hot phosgenation in the presence of a solvent or diluent, passed continuously from the bottom upward through a vertical or slanted, heated tube. After the reactants have passed through 25 this first reaction tube, they can, if appropriate, pass through a second, likewise vertical reaction tower with addition of further phosgene to complete the The advantage of vertical reaction towers reaction. which are filled with Raschig rings or other packing 30 elements to prevent rapid degassing is that the rate of isocyanate formation is increased as a result of the increased phosqene concentration at the entry point for located at the bottom due reactants the hydrostatic pressure of the column of liquid. 35 Dichlorobenzene is named as solvent. The process is a two-stage process with a cold phosgenation as first

stage and a hot phosgenation as second stage. The cold phosgenation is carried out at 0°C and the hot phosgenation is carried out with a rising temperature profile from 120°C to 160°C .

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Disadvantages of this process are the fundamental weak points of cold-hot phosgenation and also the achievable pressure. Considerably higher pressures and thus phosgene concentrations in the liquid phase can be obtained by means of pressure vessels and regulating valves. A further disadvantage is the packing elements the solid the tube reactors, since used in chloride and carbamoyl intermediates formed and precipitated in the hydrochloride phosgenation can easily lead to blockages and thus to low availability of the plant.

describes process for preparing DE 2058032 а isocyanates from amines and phosgene in the presence of an inert solvent, in which the reaction mixture 0°C in the cold phosgenation is treated at subsequently reacted fully at up to about 200°C in the hot phosgenation, with the reaction mixture from the in the hot phosgenation being conveyed phosgenation stage through a horizontal reaction zone with continual mechanical mixing and a slowly rising temperature profile. As heatable reaction vessel for the hot phosgenation, a horizontal tube through which a shaft provided with tube extends devices which longitudinal direction and has separately heatable sections of wall heating is claimed. A natural convection vaporizer (thermosyphon) can be located at the end of the tube reactor. An arrangement of three reaction tubes connected to one another in the shape of a U (system of communicating tubes) is also described. The cold phosgenation is carried out in the first, vertical tube, the hot phosgenation is carried out in

the horizontal tube and the degassing, i.e. the removal of hydrogen chloride and phosgene from the reaction mixture, is carried out in the second vertical tube. Transport pumps and regulating valves can be omitted in such a system, since the product flow is established according to the inflow rate.

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DE 2747524 describes a continuous process for preparing aromatic isocyanates in which heat is introduced into the reactor at a rate so that added phosgene does not lead to cooling and thus to caking of the carbamoyl chloride intermediate on the reactor wall. A plug flow reactor comprising two coaxial tubes into which the two amine and phosgene in an inert reactants solvent are introduced in countercurrent, each isolated and are mixed at the end of the from the other, internal tube is described. Backmixing into the feed zone is said to be prevented so as to minimize byproduct formation. Heating is by means of a steam jacket in order to prevent blocking of the mixing zone by the carbamoyl chloride intermediate. Temperatures of in general, to be necesssary; 90-140°C said are temperatures of 90-200°C are indicated. The initial 60-90°C. Practical however, temperature is, determine the upper limit to the considerations temperature. 2 atmospheres gauge is indicated as a The amine concentration in the convenient pressure. inert solvent is given as from 2 to 20%, preferably from 5 to 10%. Dichlorobenzene is preferred as inert solvent.

A tube reactor is also the preferred apparatus in the process described in WO 96/16028 for the preparation of isocyanates using isocyanate as solvent. WO 96/16028 describes a continuous, single-stage process in which the primary amine, optionally dissolved in an inert, organic solvent, is reacted with phosgene dissolved in

the isocyanate in a concentration of 10-60% by weight, based on the isocyanate/phosgene solution, at 60-180°C and pressures of 1-30 bar to form the corresponding isocyanate, with the molar ratio of phosgene to amine used being from 4:1 to 1:1 and the isocyanate used as solvent being free of solids and having a hydrolyzable chlorine content of less than 2%.

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DE 19817691 describes a two-stage process for preparing mixtures of diphenylmethane diisocyanates (MDI) 10 (PMDI) polyphenylene-polymethylene polyisocyanates having a reduced content of chlorinated by-products and a reduced iodine color number by two-stage reaction of the corresponding mixtures of diphenylmethanediamines (MDA) and polyphenylene-polymethylene-diamines 15 with phosgene in the presence of at least one organic solvent at elevated temperature, removal of the excess phosgene and solvent after the phosgenation is complete and thermal treatment of the reaction product with the molar ratio of phosgene to hydrogen chloride in the 20 residence apparatus of the second stage phosgenation being 10 - 30:1 in the liquid phase and at the same time 1 - 10:1 in the gas phase. The carbamoyl chlorides and amine hydrochlorides formed in the first stage of the phosgenation, viz. a static mixer, pass 25 through a residence apparatus in the second stage of the phosgenation in which the amine hydrochlorides are phosgenated to the corresponding carbamoyl chlorides and the carbamoyl chlorides are dissociated into the corresponding isocyanates and hydrogen chloride. 30 temperature of the first stage is usually from 40 to 60 to 130°C, particularly preferably from preferably from 90 to 120°C. Static mixers employed for in particular, nozzles. stage are, first residence apparatus for the second stage, use is made 35 of stirred apparatuses, cascades of stirred vessels and particularly preferably a column, in particular

reaction column usually having <10 theoretical plates. It is particularly advantageous to operate this column in countercurrent. The temperature at the bottom of the column is preferably from 80 to 120°C, particularly preferably from 90 to 110°C. The pressure at the top of the column is preferably from 1.0 to 4.7 atm (gauge), particularly preferably from 2.0 to 3.7 atm (gauge).

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A disadvantage of this process is that the amine hydrochloride phosgenation and the carbamoyl chloride decomposition are carried out in one and the same reactor, which leads to prolonged residence times and higher phosgene holdups.

- US 3544611 describes a process for preparing organic 15 isocyanates at a high pressure of from 10 to 50 bar. It was surprisingly found that carrying out the reaction at relatively high pressures of at least 10 atm gauge leads to higher yields of isocyanate. Furthermore, higher pressures aid the hydrogen chloride/phosgene 20 separation. The first reaction step of the isocyanate preparation, viz. the reaction between amine phosgene to form the carbamoyl chloride intermediate, is carried out in a loop reactor (mixing circuit). The second reaction step, viz. the decomposition of the 25 carbamoyl chloride to form the isocyanate, occurs in a located downstream of the reaction column hydrogen chloride-phosgene mixture is circuit. this column. Phosgene obtained at the top of condensed in two stages from this mixture. The phosgene 30 which has been condensed out is recirculated to the top of the column. At a liquid offtake in the enrichment section of the column, phosgene is taken off and recirculated to the reaction (the mixing circuit).
 - The remaining phospene is separated off from the reaction mixture taken off at the bottom of the

reaction column in a further column. In the latter, phosgene is taken off at the top, condensed in two stages in a manner analogous to the first column and recirculated to the reaction in the mixing circuit. The reaction to give the isocyanate is finished in the reaction column.

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DE 3736988 describes a continuous process for preparing organic monoisocyanates or polyisocyanates in a singlestage reaction by reacting the amine dissolved in an organic solvent with phosgene dissolved in an organic solvent in a reaction column at below 150°C. pass continuously is allowed to reaction mixture through the reaction column from the bottom upward. The reaction column has at least 10 chambers separated from one another by perforated plates. The concentration of the amine in the inert solvent is 5-40% by weight, preferably 7-20% by weight. Preferred solvents are chlorobenzene or dichlorobenzene or mixtures thereof. Phosqene is used as a 30-65% strength by weight, preferably 40-65% strength by weight, solution in the The equivalence ratio of inert solvent. phosgene is from 1:1.5 to 1:7, preferably from 1:2 to The temperature at the top of the column preferably 70-130°C, particularly preferably 90-125°C, and not more than 150°C. The mean residence time in the than 120 minutes, reaction column is not more preferably not more than 60 minutes. The pressure in the column is 1.2-3 bar abs, preferably 1.5-2.5 bar abs.

DE 3744001 likewise proposes a perforated plate column having > 10 perforated plates, preferably 20-50 perforated plates, as residence apparatus through which the reaction mixture flows from the bottom upward at a liquid velocity of 0.05-0.4 m/s, preferably 0.1-0.4 m/s, and a gas velocity of 2-20 m/s, preferably

3.5-10 m/s, and a residence time of not more than 120 minutes, preferably not more than 60 minutes. The horizontally installed perforated plates form 10-50 chambers. The temperature at the top of the reaction column is less than 150°C, preferably 70-130°C, particularly preferably 90-125°C. The pressure at the top of the column is 1.2-3 bar (abs.), preferably 1.5-2.5 bar (abs.). A mixing nozzle is claimed for the first phosgenation stage.

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EP 0291819 describes a two-stage process for preparing isocyanates by cold-hot phosgenation, in which a mixer having a rotary disk is used for the cold phosgenation and phosgenation towers are preferably used for the hot phosgenation. The phosgenation towers are operated at atmospheric pressure or a slightly superatmospheric pressure of up to 1.5 atm gauge. It is particularly advantageous to allow the reaction mixture from the cold phosgenation leaving the mixing apparatus to enter a heatable tower continuously at the top or the bottom and complete the reaction by introduction of heat. To set a particular temperature profile, a plurality of towers can also be connected in series or a combination of towers and vessels can be used.

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organic isocyanates are (US 3829458), DE 2112181 prepared continuously from primary organic amines and phosgene in an inert organic solvent in one or more reaction vessels which contain packing and through which the reactants flow in cocurrent in a transition stream. The transition stream consists of an amineliquid organic phase and a phosqenecontaining, reaction takes place containing gas phase. The reaction temperatures in the range from 50 to 220°C. In the case of incomplete reaction, the reaction mixture is circulated a number of times through the packed column. A disadvantage of this process is the high

susceptibility of the packed column to blockages caused by solids such as carbamoyl chloride, amine hydrochloride, ureas, etc., being deposited on the packing elements. Furthermore, packed columns have a high pressure drop, which results in high temperatures at the bottom and thus high thermal stress on the reaction mixture and the isocyanate formed, leading to increased by-product formation and a reduced yield.

In many processes, the reaction of phosgene and amine 10 is carried out in a loop reactor or circulating reactor in which not only the feed streams of amine and phosgene, if appropriate in a solvent, but also at least part of the reaction mixture are recycled. This dilution by recirculation of the reaction mixture 15 formed serves mainly to control the temperature and to heat so as to set achieve better removal of temperatures. The reaction between amines and phosgene is strongly exothermic. In the cases of unfavorable reaction conditions and configuration of the apparatus, 20 increased by-product temperatures cause formation which, for example in the case of tolylene diisocyanate (TDI), leads to a decrease in yield and formation of tar. Main by-products formed are ureas.

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(BASF) describes a mixing circuit process DE 2624285 for the continuous preparation of organic isocyanates from organic amines and phosgene in the presence of organic solvents, in which the phosgene is mixed into the circulated reaction solution and the reaction mixture obtained and the amines or amine solution are fed into the mixing and reaction zone in such a way that an energy dissipation density of from 5 to 1000 kJ per m³ of recirculated reaction mixture plus introduced amine solution is produced therein. The reaction is carried out at from 90 to 220°C, preferably from 120 to in a pressure range from 1 to 10 bar, and 180°C,

preferably from 1 to 3 bar. The residence times are from 10 to 180 minutes. The molar ratio of amine to phosgene is such that from 1 to 10 mol, preferably from 1.3 to 4 mol, of phosgene per amino group are present in the reaction mixture. The yields are from 88 to 98% by weight, based on the amine used.

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The mixing circuit process described in DE 2624285 is developed further in EP 0150435. In the process for the preparation of organic isocyanates continuous reaction of organic amines with phosgene the presence of organic solvents, with hydrogen chloride being separated off and the reaction mixture being partly circulated, the hydrogen chloride content of the reaction mixture recirculated to the addition of amine after the hydrogen chloride has been separated off is, prior to the addition of amine, equal to or less than 0.5% by weight, preferably from 0.01 to 0.4% by weight, based on the total weight of the reaction mixture, and the molar ratio of phosgene to amino groups of the 20 organic amines is 12 - 200:1. The reaction is carried 100-220°C, preferably 120-180°C, and in at pressure range of 5-100 bar, preferably 15-50 bar.

a process for the continuous DE 3403204 describes 25 organic isocyanates, preferably of preparation of organic amines, polyisocyanates, reaction by preferably polyamines, with phosgene in the presence of organic solvents under superatmospheric pressure, e.g. from 5 to 100 bar, and elevated temperatures, e.g. from 30 100 to 220°C, with the reaction mixture being partly by the natural convection circulated, preferably principle, and the hydrogen chloride content of the reaction mixture prior to the addition of amine being less than 0.5% by weight, based on the total weight of 35 the reaction mixture, and the molar ratio of phosgene to amino groups of the organic amines being 12 - 200:1.

DE 3212510 describes a process for the continuous preparation of organic isocyanates. The primary organic amine is brought into contact in a virtually dispersed state with an excess of phosgene at a gauge pressure of 5 10 kg/cm², approximately 10 bar, and a temperature of formation of 100°C, resulting in 60 to corresponding organic carbamoyl chloride from organic amine and hydrochloride formed as intermediate. Hydrogen chloride is formed as by-product. In this 10 first stage of the reaction, from 30 to 70% of the carbamoyl chloride is converted into isocyanate. reaction mixture is maintained at a gauge pressure of 10 kg/cm² and a temperature of from 120 to 160°C, that the hydrochloride is converted into carbamoyl 15 chloride and the conversion of the carbamoyl chloride into isocyanate is complete. The reaction takes place in a circulation reactor (circulation line) or in a tank-shaped reaction vessel. In the first case, the phosgene is allowed to circulate together with the 20 solvent in a tubular circulation line and the amine is mixed in in this (mixing circuit). The residence time in the first stage is 30-120 minutes and that in the second stage is 10-120 minutes. ortho-Dichlorobenzene is chosen as solvent. 25

likewise describe loop and DE 1811609 GB 763535 reactors or circulation reactors (mixing circuits as reaction system). The organic isocyanate is prepared by reacting an amine with phosgene in a single-stage continuous reaction with circulation of isocyanate, solvent and unreacted phosgene. The pressure described as sufficient in the process described in GB 763535 is 5-20 pounds per square inch, the reaction temperature is 90-180°C, the TDA concentration in the solvent is 5-30%, the stoichiometric excess of phosgene is at least 25%, preferably 70-110%, and the solvents used are

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hydrocarbons, preferably chlorinated aromatic dichlorobenzene. In DE 1811609, the organic amine, ortho-dichlorobenzene or appropriate in solvent, and an excess of phosgene are mixed under high shear stress into the circulating reaction mixture, by which means conditions which advantageously deviate from GB 763535 can be set owing to the mixing. The reaction pressure is preferably at least 1.8-14·10⁵ Pa, preferably 4.2·10⁵ Pa $3.5 \cdot 10^{5} \text{ Pa.}$ orThe temperature is stated to be preferably 102-130°C and, in the case of toluenediamine, preferably 90-120°C. The excess of phosgene is 50-200%, preferably 70%.

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DE 1037444 (US 2822373) describes a continuous process for preparing organic isocyanates, in which a solution 15 of the organic amine in an inert solvent is reacted with a solution of phosgene in an inert solvent at from 90 to 180°C in a reaction zone in which superatmospheric pressure and turbulent flow prevail. The reaction solution is then depressurized into a zone at a lower 20 pressure, usually atmospheric pressure, and hydrogen chloride and phosgene are taken off as a gaseous mixture. The isocyanate is separated from the solvent by distillation. In the process described, the amine solution is introduced into a pumped circulation line, 25 the circulating mixture is heated downstream of the reaction zone by means of a heat exchanger and is then depressurized via a throttle valve into a tank-shaped residence vessel. From this, the reaction mixture is either taken out for the pumped circuit or partly 30 discharged for removal of solvent and recovery of the isocyanate. The solvent is condensed from the gaseous hydrogen chloride, excess phosgene mixture of solvent taken off from the reservoir in a condenser and is returned to the reservoir. Preferred pressures in 35 the pumped circulation line are given as 5-20 pounds per square inch. The amine concentration in the solvent

is 5-30% by weight, and the amount of phosgene is at least 1.25 mol per amino group of the amine. The preferred solvent is stated to be orthodichlorobenzene.

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US 3574695 describes an improved continuous process for preparing organic isocyanates. The residence times can be shortened by treating the product from the first reaction zone in the second reaction zone with at least 0.75 mol of phosgene per equivalent of organic amine added in the first reaction zone. The residence time in this second reaction zone is 5-45 minutes at a temperature of at least 130°C. A gaseous mixture of hydrogen chloride and phosgene and a liquid solution of the organic isocyanate in the solvent are continuously taken off from the second reaction zone.

GB 1034285 describes a continuous process for preparing organic isocyanates by reaction of phosgene with a primary polyamine in the presence of an inert organic solvent, with the reactants being fed separately from one another into a tube reactor and brought contact there and a mixture of the same solvent, the being recirculated mixture and phosgene reaction through this tube reactor. As reactor, it is possible to use an assembly of two cylindrical vessels between which the reaction mixture is circulated or an annular tube reactor. The reaction mixture can be stirred by means of stirrers. The temperature in the tube reactor The pressure is atmospheric pressure is $8-50^{\circ}C$. above this. The concentration slightly introduced primary polyamine in the solvent is 2-20% by weight. The amount of phosgene added to the stream circulated by pumping is from 5 to 20 mol of phosgene per mol of amino groups in the polyamine solution inert organic solvent, use is made As chlorobenzene or ortho-dichlorobenzene.

GB 1212249 describes a process for preparing isocyanates in which an excess of phosgene is reacted with a forward-directed, mixed film of the amine in an inert solvent in the first stage. A cylindrical tube is regarded as suitable for producing this film.

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JP 57-048954 describes a process for preparing organic isocyanates in which the solution of a primary amine is introduced just upstream of the feed point of a static mixer which is located in a circulation reactor. A solution of phosgene in an organic isocyanate circulates in the circulation reactor.

- 15 Phosgenation in a loop reactor is also described in JP 60-10774 in which an isocyanate-containing reaction mixture is circulated by pumping. However, high yields are only achieved at amine concentrations of 5-10%.
- The low temperatures in the first stage and the high 20 temperatures in the second stage of loop reactor or are energetically processes circuit mixing disadvantageous. Since the reaction between an organic amine and phosgene is strongly exothermic, intensive cooling has to be employed in the first step in order 25 maintain the desired reaction temperature. viz. the decomposition the reaction, second form the isocyanate, carbamoyl chloride to significantly endothermic, so that the reaction mixture has to be heated again in the second stage. 30

Moreover, the significantly lower chemical yield compared to processes carried out in a single pass is particularly disadvantageous, since isocyanate already formed reacts with amine to form ureas in the mixing circuit due to backmixing. To suppress this secondary reaction, a low maximum steady-state isocyanate

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DE 102 60 082 A1 2004.07.01

concentration is frequently permitted, but this in turn results in low space-time yields.

EP 0716079 for the described in the process continuous preparation of organic isocyanates, the reaction mixture is circulated at 60-100°C in a bubble process described is carried out The column. slightly subatmospheric or slightly superatmospheric pressure, in general at 0.5-5 bar, preferably 1-3 bar.

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EP 0570799 describes a process in which the reaction and phosgene to form isocyanate between amine carried out in the gas phase. Gas-phase phosgenation is known for the preparation of aliphatic diisocyanates diisocyanates (EP 0570799), aromatic (EP 0289840), cyclic diisocyanates (EP 1078918) and of triisocyanates EP 0749958, EP 0676392 and EP 0289840 (EP 0749958). for preparing aliphatic describe processes gas-phase and triisocyanates by diisocyanates phosgenation in which the reactants are mixed between nozzle and tube on entering the tube reactor described through nozzles or a combination of nozzles and an annular gap. A Reynolds number of RE > 4700 in the tube is indicated as an essential criterion for mixing here. EP 0570799 in mixer is proposed aromatic diisocyanates gas-phase by preparation of phosgenation.

DE 1192641 describes a process for preparing primary aromatic of by reaction 30 isocyanates araliphatic amines with phosgene in the presence heating of the subsequent solvents and mixture, with the isocyanate which is to be prepared in the reaction being used as solvent.

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DE 100 27 779 claims a process for preparing isocyanates by reacting amines with phosgene, in which

the isocyanate is used as solvent and the reaction is carried out in a reaction column and the condensed phase at the bottom of the reaction column is recirculated in its entirety or in part to the enrichment section of the reaction column. The number of theoretical plates in the reaction column is 5-60. The temperature is from -20°C to 300°C and the absolute pressure is 0.2-60 bar.

US 2683160 describes a process for preparing aromatic 10 isocyanates in which gaseous phosgene and a solution of an aromatic amine in a chlorinated aromatic hydrocarbon introduced simultaneously into solvent are as isocyanate in the the desired solution of abovementioned solvent. The solution of the desired 15 isocyanate in the solvent is maintained at the boiling point of the solvent, i.e. at 130°C - 300°C, with the solution boiling under reflux. The by-product hydrogen chloride and excess, unreacted phosgene are taken off continuously through the reflux condenser. The excess 20 of phosgene is at least 50%, preferably from 80 to The stoichiometric amount. of the 5 is from to 30%, concentration in the solvent preferably from 8 to 12%. The reaction is carried out in a single apparatus which is stirred well and heated 25 sufficiently.

DE 2252068 describes a process for preparing organic isocyanates from amine and phosgene, in which the amine which has been preheated to a temperature below its decomposition temperature at superatmospheric pressure is firstly reacted with preheated phosgene in the presence of an excess of an organic isocyanate as solvent at temperatures and pressures such that the reaction proceeds in a homogeneous, liquid phase and the organic carbamoyl chloride formed as intermediate is subsequently thermally dissociated at a lower

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pressure in a second stage. In a preferred embodiment, the first reaction stage is carried out adiabatically. The reaction components are fed in at temperatures in the range 120-180°C. The temperature of the reaction mixture at the outlet is maintained at 180-250°C and the pressure is maintained at 100-300 atm. The residence time of the components in the first reaction zone should be 5-150 seconds. The second reaction stage is carried out isothermally. The feed temperature is 120-250°C and the pressure is 3-30 atm. The residence time The isocyanate taken off from the is 3-30 minutes. 50-80°C prior to is cooled to second stage recirculation.

US 3801518 describes a process for preparing tolylene diisocyanate having an increased acidity of 0.03-0.3% by weight. This is achieved by phosgenation of toluenediamine and subsequent residence of the reaction product in a phosgene atmosphere at 100-200°C for a period of at least 0.08 hour, preferably from 0.08 hour to 2 hours.

US 3912600 describes the reduction in the acidity and the content of hydrolyzable chlorine in a polymethylene-polyphenylene polyisocyanate (PMDI) by treatment of this in an inert, organic solvent at a pressure of 20-60 psia and a temperature of 150-230°C, with removal of volatile compounds, known as low boilers.

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GB 1196008 describes a continuous process for preparing aromatic monoisocyanates or diisocyanates by phosgenation of the corresponding amines in an organic solvent at 120-200°C in two reaction vessels coupled to one another, with the excess of phosgene over the calculated stoichiometric amount being 5-20%.

It is an object of the present invention to develop a two-stage or multistage process which gives isocyanates in very high chemical yields and with high space-time yields at a low phospene holdup.

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surprisingly been found that, contrary to has generally prevailing opinion, the second reaction, viz. the phosgenation of the amine hydrochloride, proceeds at a high reaction rate at high phosgene concentrations and elevated temperatures. High pressures are therefore 10 advantageous for this reaction, since high pressures imply high phosgene concentrations in the liquid phase. Furthermore, elevated temperatures are advantageous for achieving high space-time yields. I.I. Konstantinov, A.I. Kormucheshkina, Zhurnal Prikladnoi Khimii, 49 (3), 15 pp. 596-599, 1976) state that the phosgenation of the amine hydrochloride is very slow and is the ratedetermining step of the overall reaction cycle to the al. present Konstantinov et isocyanate. measurements and quantify the reaction rates. According 20 to them, the reaction rate for the phosgenation of the hydrochloride is considerably lower than that for the free amine. As described in GB 1212249, the formation hydrochloride also leads to a loss amine isocyanate yield due to urea formation. Since 25 solubility of amine hydrochlorides in the corresponding commercially also in most reaction mixtures and available solvents is very low, hydrochloride formation drastically increases the problem of solids also

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DE 3323882 describes a continuous process for the hot phosgenation of amine hydrochloride or mixtures thereof with carbamoyl chloride suspended in solvents by means of excess phosgene at from 80°C to 200°C, preferably from 100°C to 180°C. In this process, the solids are retained in the reactor by means of a suitable

separation device and the isocyanate which is formed during the reaction and is present as a solution in the solvent is continuously taken off from the reactor. The solids are preferably separated off by means of a filter. Disadvantages of this process for phosgenation of hydrochlorides are the complicated solids handling, the risk of blockage of pipes and, in particular, of regulating valves and flowmeters and also the long residence time which requires large apparatuses and results in a high phosgene holdup and also the severe reaction conditions and the relatively low yields.

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preparing process for describes а DE 2404773 monoisocyanates, diisocyanates and/or polyisocyanates from organic primary amines and phosgene, in which the 15 primary amines are mixed with at least 3 phosgene per amino group in the absence of a solvent and the reaction mixture is simultaneously broken up to a mean particle size of 1-100 μm and the resulting carbamoyl chloride and of suspension 20 converted into the is hydrochloride in phosgene from 100 to isocyanates at corresponding preferably from 120 to 160°C, and pressures of from 14 to 55 bar, preferably from 21 to 41 bar. The process is a two-stage process in which the starting materials 25 primary amine and phosgene are mixed in the first stage at from -30°C to 60°C, preferably 0-50°C, at atmospheric pressure or preferably superatmospheric pressure, particular at from 14 to 55 bar, and at the same time the particles are comminuted to a mean particle size of 30 from 1 to 100 μm , preferably from 1 to 50 μm . The amine is added as a liquid, melt or possibly as a powder to the phosgene. Various mixing and comminution devices are described. The second stage comprises the reaction of amine hydrochloride with phosgene to form carbamoyl 35 its decomposition into isocyanate chloride and hydrogen chloride in a pressure vessel at 100-180°C,

preferably from 120 to 160°C, and pressures of 14-55 bar, preferably from 21 to 41 bar. This process is technically very complicated and not economical.

5 DE-A 156844 likewise describes a phosgenation of an amine hydrochloride suspension which is carried out at elevated temperature in a multistage cascade of stirred vessels. A particular disadvantage of a cascade of stirred vessels is the high phosgene holdup.

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It has now surprisingly been found that the phosgenation of the amine hydrochloride is a fast reaction at high phosgene concentrations and elevated temperatures.

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Furthermore, it has surprsingly been found that amine hydrochloride and carbamoyl chloride formed in situ display a high level of supersaturation in the reaction mixture when a static mixer, preferably a nozzle, used as reactor for the first stage. Even when amine 20 hydrochloride and/or carbamoyl chloride precipitate, the process claimed has considerable advantages since a hydrochloride particle size narrow amine distribution a very small mean having diameter, usually in the nanometer to micron range, can 25 be produced when using a static mixer, preferably a nozzle, as reactor for the first stage as a result of the introduction of high mixing energies. However, is more advantageous to achieve high conversions or if possible complete conversion in the phosgenation of 30 amine hydrochloride before solid amine hydrochloride or carbamoyl chloride precipitates in relatively large amounts, since, firstly, solids handling is complicated from a process engineering point of view and can lead and blockages and, secondly, to caked material 35 large and agglomerated phosgenation οf hydrochloride particles as described in the literature

is very slow.

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The present invention accordingly provides a process for preparing polyisocyanates by reacting organic amines with phospene, wherein the reaction is carried out in at least three stages, with the first stage being carried out in a mixing apparatus, the second stage in at least one residence apparatus and the third stage in at least one separation apparatus and the pressure in each successive stage being lower than that in the previous stage.

The first stage of the process of the present invention comprises essentially the reaction of the amine carbamoyl chloride and amine hydrochloride, the second stage comprises essentially the reaction of the amine formed in the first stage to hydrochloride third stage comprises chloride and the carbamoyl essentially the dissociation of the carbamoyl chloride into isocyanate and hydrogen chloride.

In the process of the present invention, the reaction between organic amine and phosgene is carried out in three or more stages in an inert solvent, preferably toluene or chlorobenzene, dichlorobenzene or mixtures thereof, using an excess of phosgene, with the pressure stage to stage. The reduced from phosgenation stage comprises a static mixer, preferably a nozzle. The pressure upstream of the nozzle preferably from 3 to 70 bar, in particular from 15 to 45 bar. The pressure difference over the nozzle is at least 0.5 bar. The temperature in the first stage is preferably from 80 to 190°C, in particular from 90 to 150°C. The second stage comprises one or more residence apparatuses, preferably one residence apparatus, which is operated at a pressure of from 2.5 to 35 bar, preferably from 15 to 35 bar. Downstream of the nozzle,

the reaction mixture is depressurized by means of a valve or some other device suitable for this purpose to the pressure of the residence apparatus of the second stage. However, the natural pressure drop of the nozzle can also be used for the pressure reduction.

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The reactor of the first stage can also be integrated into the reactor of the second stage. In particular, a mixing nozzle can dip into the gas phase or preferably into the liquid phase of the second reactor, i.e. can be located entirely or partly therein. It is also possible for the output from the nozzle to be conveyed by means of a pipe, an immersed tube or a plug-in tube into the gas phase or preferably into the liquid phase of the reactor of the second stage.

The temperature in the second stage is from 80 to 190°C, preferably from 90 to 150°C. Possible types of reactor tube reactors, the second stage are apparatuses, unstirred residence 20 other apparatuses. apparatuses and separation reactor can also be provided with a pumped circuit which may in turn have a heat exchanger for setting the reactor temperature. In the case of a stirred vessel, an unstirred residence apparatus or possibly also in 25 the case of a phase separation apparatus, the liquid phase is preferably depressurized under level control gas phase is depressurized under pressure and the control into the reactor of the third stage. However, which comprises mainly phosgene, phase, 30 the qas hydrogen chloride and possibly solvent, can also be passed directly to the work-up, e.g. fractionation into hydrogen chloride and solvent phosgene, mixtures thereof. The residence reactor of the second stage can, depending on the desired residence time and 35 capacity of the plant, have relatively large dimensions and volumes, which can be regarded as disadvantageous

from the point of view of cost or safety, e.g. phosgene holdup at high pressure. In this case, the reactor of the second stage can also be realized as two or more similar or different reactors and types of reactor, which can be connected in parallel or, if appropriate to influence the residence time spectrum, also in series.

The reactor of the third stage of the process of the present invention is operated at a pressure of from 2 to 20 bar, preferably from 3.5 to 16 bar. Downstream of the residence reactor of the second stage, the reaction mixture is depressurized to the pressure of the third reactor by means of a valve or some other device suitable for this purpose. A natural pressure drop may also be able to be utilized.

In any case, the pressure in the following stage is, as described above, selected so that it is lower than in the previous stage.

The temperature in the third stage is from 80 to 190°C. The type of reactor used for the third stage is a а reaction column particular column. in described, for example, in WO 99/54289. The temperature at the bottom is from 80 to 190°C and the temperature at the top is from 50 to 120°C. The column used as reactor of the third stage can also be utilized for removing the excess phosgene from the reaction mixture. reactor of the third stage can, like the reactor of the second stage, be disadvantageously large. In this case, the reactor of the third stage can also be realized as two or more similar or different columns connected in series.

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The output from the bottom of the reaction column is worked up by customary methods to remove any phosgene

still present and to separate off the solvent. In the case of the preparation of TDI, the crude TDI is subsequently subjected to removal of high boilers and purification by distillation. Phosgene, hydrogen chloride and possibly solvent can be separated off in a known manner from the vapor leaving the reaction column and, if appropriate, the residence reactor of the second stage and can, if appropriate, be recirculated.

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As solvents, preference is given to using chlorinated 10 hydrocarbons dichlorobenzene, aromatic such as chlorobenzene, trichlorobenzene or mixtures thereof, aromatic or aliphatic hydrocarbons such as toluene, hexane, heptane, pentane, xylene, benzene, biphenyl, ketones such as 2-butanone, cyclohexane, 15 such as isobutyl ketone, esters methyl ethyl acetate, butyl acetate, nitriles isophthalate, such as acetonitrile and also sulfolane.

The particular advantage of the process of the present 20 is that, in contrast to the invention customary in the prior art, the two reaction steps 1) phosqenation the amine hydrochloride form οf decomposition of the carbamoyl chloride and 2) chloride isocyanate and hydrogen into 25 carbamoyl chloride are carried out partly or entirely in separate stages or reactors and very high chemical yields, very high space-time yields and at the same time a very low phosgene holdup can be achieved as a result of the independent setting of the optimum pressure and optimum 30 and respective reaction for the temperature selection of the most favorable reactor design in each case. The synthesis can be carried out adiabatically or isothermally. The differing apparatus design optimally takes account of the conditions of the two reactions. 35 While the phosgenation of the amine hydrochloride requires high pressures, low pressures are advantageous

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decomposition οf the carbamoyl chloride. for the times selected for the the residence Furthermore, reactor for the phosgenation of the amine hydrochloride can be shorter than those for the decomposition of the which considerably carbamoyl chloride, reduces removal, overall phosgene holdup. Furthermore, of the gaseous hydrogen stripping, particular by chloride formed is advantageous for the decomposition carbamoyl chloride because the carbamoyl the chloride/isocyanate equilibrium is in this advantageously shifted to the side of the isocyanate. This can be taken into account by the (reaction) column as optimum reactor of a choice design. The excess phosgene can also be removed at the same time. This is not absolutely necessary in this stage, and can also be carried out in a further stage. In contrast, the removal of hydrogen chloride in the residence reactor of the second stage would be very disadvantageous, since the phosgene necessary for the phosgenation of the amine hydrochloride would in such a removed together with the hydrogen also be chloride. Vaporizing phosgene also leads to cooling of reaction mixture, which could lead to massive precipitation of solid carbamoyl chloride and amine hydrochloride.

The rapid reaction between amine and phosgene to form carbamoyl chloride and hydrogen chloride and also amine requires high pressures as hydrochloride conditions in both the first and second stages achieve high phosgene concentrations in the phase and thus high excesses of phosgene if chemical yields are to be obtained as a result of low In addition, good mixing by-product formation. necessary, i.e. a static mixer, preferably a nozzle, should be used as apparatus. High admission pressures upstream of the nozzle allow high pressure drops over

the nozzle and thus the introduction of high mixing energies.

Dissolved amine hydrochloride and very small amine hydrochloride particles surprisingly react very quickly with phosgene in the residence reactor of the second stage and therefore do not require a long residence time. High phosgene concentrations are advantageous here. Relatively high pressures do not have an adverse effect on the phosgenation of the amine hydrochloride, 10 so that high phosgene concentrations in the liquid phase can preferably be set by means of relatively high pressures. Suitable apparatuses are, in particular, tube reactors, stirred vessels, unstirred residence apparatuses, phase separation apparatuses or other 15 Furthermore, apparatuses. residence embodiments of these apparatuses are advantageous in order to compensate, if appropriate, temperature drops decomposition the endothermic caused by carbamoyl chloride. The reaction of the third stage, 20 viz. the decomposition of the carbamoyl chloride into hydrogen chloride, is а pressureisocyanate and dependent equilibrium reaction. It is advantageously shifted to the side of the desired isocyanate by low pressures. Since this reaction does not require any 25 phosgene, the low phosgene concentrations in the liquid phase characteristic of low pressures do not interfere and in fact lead to a low phosgene holdup in the reactor of the third stage, which is advantageous from a safety point of view. In this way, the overall 30 phosgene holdup in the process, and possibly also in apparatuses, can be significantly individual reduced compared to a cascade of stirred vessels or a reaction tower. High pressures are, in contrast, very unfavorable for the dissociation of carbamoyl chloride 35 and require long residence times, high temperatures and energy consumptions, since at high pressures the

equilibrium lies far on the side of the carbamoyl chloride. Long residence times in turn result in a high phosgene holdup. A (reaction) column as described, for example, in WO 99/54289 (DE 19817691) is particularly useful as apparatus for the third stage. In addition the stripping effect shifts the carbamoyl chloride/isocyanate equilibrium advantageously to the side of the desired isocyanate.

The phosgenation of the amine hydrochloride does not 10 have to be completed in the second stage, and it is the decomposition of likewise possible for the chloride to the second stage. start in carbamoyl However, preference is given to a design of the reactor of the second stage in terms of residence time and 15 other process engineering parameters which is such that hydrochloride is phosgenation of the amine decomposition the virtually complete and carbamoyl chloride has not yet progressed to any large extent. 20

If the phosgenation of the amine hydrochloride and the carbamoyl chloride are, decomposition of the described in the prior art, carried out in one stage or in one reactor, the high pressure required for the phosgenation of the amine hydrochloride results in a the carbamoyl chloride into the low conversion of thus long residence times. A isocyanate and phosgene concentration and long residence times (large in turn result in a very high reaction volumes) phosgene holdup. This is also present at high pressures and temperatures which are of concern from a safety separation Physical of the of view. point amine of the viz. the phosgenation reactions, hydrochloride in the second phosgenation stage at a high pressure and the decomposition of the carbamoyl chloride in the third phosgenation stage at a low

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pressure, enables high chemical yields, high space-time yields and especially a low phosgene holdup in the overall process and possibly also in the individual apparatuses to be achieved.

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Physical separation of the first and second stages is not absolutely necessary, since a high pressure increases the phosgene concentration in the liquid phase, which favors both the first reaction between amine and phosgene and the second reaction between amine hydrochloride and phosgene.

Furthermore, the process can be carried out at elevated temperature and if appropriate also isothermally in all stages. This results, in particular, in high space-time low phosgene holdups and yields and thus together simultaneously higher with apparatuses chemical yields, especially compared to the classical cold/hot phosgenation. In addition, avoiding cooling of the reaction mixture in the first stage and subsequent reheating in the second stage and the following stages saves a considerable quantity of energy. Avoiding the precipitation of amine hydrochloride as solid enables long residence times, which can sometimes be realized only by means of a circulation mode of operation (loop reactors), to be avoided. Although the circulation mode of operation likewise involves a lower phosgene holdup than, for example, a cascade of stirred vessels, suffers from increased formation of by-products, particular ureas. To avoid urea formation, the amine or isocyanate concentration has to be kept low, which leads to very low space-time yields.

The temperatures and pressures employed are to some some extent dependent on the amine used. Likewise, the phosgene excesses and residence times to be employed in the individual apparatuses are dependent on the amine

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the case of diphenylmethane diisocyanates used. polyphenylene-polymethylene and/or (MDI) polyisocyanates (PMDI) or mixtures of these two, the excess of phosgene should be at least 100% of the stoichiometric amount, while in the case of tolylene diisocyanate (TDI), it should at least 300% of the stoichiometric amount and in the case of hexamethylene diisocyanate (HDI) and isophorone diisocyanate (IPDI) 300% of least likewise be at it should stoichiometric amount. The residence time in the first stage (static mixer) is naturally very short and is defined by the apparatus design. The mean residence time in the residence apparatus of the second stage can be from one second to 30 minutes. Preference is given residence times of from 30 seconds mean particularly preferably from 2 to 10 minutes, minutes. The mean residence time in the apparatus of the third stage (reaction column) is dependent on the number of theoretical plates, the height of the weir, the volume of liquid phase, the throughput through the 20 column and other process engineering parameters. It is preferably not more than 1 hour.

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concentration in the inert solvent is amine The amine used and on on the 25 likewise dependent In the case of diphenylmethanediamine (MDA) solvent. and/or polyphenylene-polymethylenepolyamines (PMDA) mixtures of these two, it can be 5-50% by weight, case in the 25-40% by weight; preferably 5-50% by weight, (TDA) it can be toluenediamine 30 by weight; in the 15-30% preferably hexamethylenediamine (HDA) it can be 5-50% by weight, the case 15-30% by weight; and in preferably isophorone diisocyanate (IPDI) it can likewise be 5-50% by weight, preferably 15-30% by weight. The phosgene 35 concentration in the inert solvent can be 0-70% by weight, preferably 10-50% by weight. Preference

given to using the same solvent as for the amine. It is also possible to omit a solvent entirely.

The invention is illustrated by the following examples.

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Examples:

la) Tube reactor as residence apparatus (I)

A solution of 0.73 kg/h of toluenediamine 10 3.3 kg/h of chlorobenzene was reacted at 110°C with 6.2 kg/h of phosgene in a mixing nozzle. The pressure drop over the nozzle was 6 bar. The reaction mixture was depressurized directly into a tube reactor heated by means of a double wall as residence apparatus having a 15 residence time of 2 minutes at 110°C and a pressure of The output from the tube reactor was 15 bar abs. depressurized via a regulating valve into a bubble cap tray column (reaction column). The temperature at the bottom of the reaction column was 150°C and the pressure 20 at the top was 3.5 bar abs. At the top of the column, a of phosgene (4.8 kg/h,(5.8 kg/h)weight), hydrogen chloride (0.85 kg/h, 15% by weight) and chlorobenzene (0.19 kg/h, 3% by weight) and small amounts of various low boilers (CCl $_4$, CHCl $_3$, N $_2$, CO, CO $_2$) 25 taken off, partially condensed and passed separation by а hydrogen chloride/phosgene method. Part of the condensate was returned to the column as runback. The temperature at the top was 71°C. At the bottom of the column, a mixture (4.4 kg/h) of 30 diisocyanate (1.0 kg/h, 23% by weight), tolylene 70% weight), phosgene (3.1 kg/h,by chlorobenzene weight), hydrogen chloride (0.27 kg/h,6% by (0.02 kg/h, 0.5% by weight) and small amounts of high boilers (0.04 kg/h, 1% by weight) was taken off. A 35 shell-and-tube apparatus having 13 tubes was used as

bottom circulation vaporizer.

1b) Tube reactor as residence apparatus (II)

A solution of 0.73 kg/h of toluenediamine (TDA) 3.2 kg/h of chlorobenzene was reacted with 6.2 kg/h of phosgene in a mixing nozzle. The pressure drop over the reaction mixture 8 bar. The was nozzle depressurized directly into a tube reactor as residence apparatus having a residence time of 10 seconds at 120°C and a pressure of about 15 bar abs. The output from the 10 tube reactor flowed directly into a bubble cap tray column. The temperature at the bottom of the reaction column was 150°C and the pressure at the top was 15 bar abs. At the top of the column, a mixture (3.0 kg/h) of phosgene (2.1 kg/h, 71% by weight) and 15 hydrogen chloride (0.85 kg/h, 29% by weight) and small amounts of chlorobenzene and various low boilers (CCl4, $CHCl_3$, N_2 , CO, CO_2) was taken off, partially condensed and passed to hydrogen chloride/phosgene separation by a known method. Part of the condensate was returned to 20 the column as runback. At the bottom of the column, a mixture (kg/h) of tolylene diisocyanate (1.0 kg/h, 14% by weight), chlorobenzene (3.2 kg/h, 45% by weight), phosgene (2.9 kg/h, 41% by weight) and small amounts of high boilers (0.05 kg/h, 1% by weight) was taken off. A 25 shell-and-tube apparatus having 13 tubes was used as bottom circulation vaporizer.

2) Stirred vessel as residence apparatus

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A solution of 0.73 kg/h of toluenediamine (TDA) in 3.3 kg/h of chlorobenzene was reacted at 140°C with 6.2 kg/h of phosgene in a mixing nozzle. The pressure drop over the nozzle was 4 bar. The nozzle was integrated into a tube reactor as residence reactor into which the reaction mixture was depressurized. The residence reactor was a stirred vessel which was

brought to a temperature of 140°C via the double wall. The stirrer speed was 1000 revolutions per minute. As an alternative, the nozzle was placed outside the residence reactor and the output from the nozzle was fed directly into the liquid phase via a plug-in tube. Depressurization into the gas phase led to somewhat lower yields of tolylene diisocyanate (TDI). The liquid phase is discharged under level control and the gas phase is discharged under pressure control from the into a bubble cap tray column residence reactor 10 (reaction column). The stirred vessel was operated in the pressure range from 2.5 to 35 bar abs. The mean residence time of the liquid phase (regulated via the level) was up to 30 minutes. The stirred vessel was also operated with an external loop provided with a 15 heat exchanger (pumped circuit with gear pump). The temperature at the bottom of the reaction column is 110°C and the pressure at the top is 3.5 bar abs. At the top of the column, a mixture (5.0 kg/h) of phosgene hydrogen chloride weight), (4.0 kg/h)20% by 20 and chlorobenzene weight) 17% bv (0.85 kg/h)(0.16 kg/h, 3% by weight) and small amounts of various low boilers (CCl₄, CHCl₃, N₂, CO, CO₂) is taken off, hydrogen condensed and passed to partially chloride/phosgene separation by a known method. Part of 25 the condensate was returned to the column as runback. The temperature at the top was 70°C. At the bottom of tolylene mixture (5.2 kg/h)of column, a the diisocyanate (1.0 kg/h, 19% by weight), chlorobenzene (3.1 kg/h, 59% by weight), phosgene (1.15 kg/h, 22% by 30 weight), hydrogen chloride (0.02 kg/h, 0.3% by weight) and small amounts of high boilers (0.02 kg/h, 0.4% by was taken off. A shell-and-tube apparatus weight) bottom circulation was used as 13 tubes having vaporizer. 35

3) Phase separation apparatus

A solution of 0.74 kg/h of diaminodiphenylmethane (MDA) reacted with was 1.6 kg/hof chlorobenzene 2.1 kg/h of 1.9 kg/h of phosgene in solution of chlorobenzene in a mixing nozzle. The pressure drop 5 over the nozzle was 5 bar. The nozzle was depressurized via a plug-in tube into the liquid phase of a phase separator. The pressure downstream of the nozzle was 12 bar and the temperature was 115°C. The liquid phase and the gas phase were conveyed separately into a 10 bubble cap tray column (reaction column). The mean residence time of the liquid phase was about 3 minutes. The temperature at the bottom of the reaction column was 115°C and the pressure at the top was 5 bar abs. At the top of the column, a mixture (1.5 kg/h) of phosgene 15 chloride weight), hydrogen (0.73 kg/h)50% by and chlorobenzene (0.50 kg/h)33% by weight) (0.24 kg/h, 16% by weight) and small amounts of various low boilers (CCl $_4$, CHCl $_3$, N $_2$, CO, CO $_2$) was taken off, hydrogen passed to condensed and 20 partially chloride/phosgene separation by a known method. Part of the condensate was returned to the column as runback. The temperature at the top was 110°C. At the bottom of the column, a mixture (4.9 kg/h) of methylenedi(phenyl 0.93 kg/h,19% by weight), (MDI, isocyanate) 25 71% by weight), phosgene (3.5 kg/h)chlorobenzene and hydrogen chloride (0.43 kg/h, 9% by weight) (0.05 kg/h, 1.0% by weight) was taken off. A shell-andtube apparatus having 13 tubes was used as bottom circulation vaporizer. 30

We claim:

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- for preparing polyisocyanates process reacting organic amines with phosgene, wherein the reaction is carried out in at least three stages, with being carried out in a stage apparatus, the second stage in at least one residence the third in at least stage apparatus and pressure in each the apparatus and separation successive stage being lower than that in the previous stage.
- 2. A process according to claim 1, wherein the polyisocyanate is diphenylmethane diisocyanate (MDI), polyphenylene-polymethylene polyisocyanate (PMDI) or a mixture of these two, tolylene diisocyanate (TDI), hexamethylene diisocyanate (HDI) or isophorone diisocyanate (IPDI).
- 20 3. A process according to claim 1, wherein a nozzle is used as apparatus for the first stage.
- A process according to claim 1, wherein a tube reactor, a stirred vessel, an unstirred residence
 apparatus or a phase separation apparatus for gas and liquid phases is used as apparatus for the second stage.
- 5. A process according to any of claims 1-4, wherein the residence time in the residence apparatus of the second stage is from 1 second to 30 minutes, preferably from 30 seconds to 10 minutes, particularly preferably from 2 to 7 minutes.
- 35 6. A process according to any of claims 1-5, wherein the residence reactor of the second stage is configured

as two or more reactors of the same or different types which are connected in parallel or in series.

- 7. A process according to claim 1, wherein one or more (reaction) columns are used as apparatus for the third stage.
- 8. A process according to claim 1, wherein the phosgene is separated off in the apparatus of the third stage, preferably a (reaction) column.
- 9. A process according to claim 1, wherein the pressure is reduced from the pressure of the reactor of the first stage to the pressure of the reactor of the second stage by means of a regulating valve or some other device characterized by a pressure drop.
 - 10. A process according to claim 1, wherein the pressure is reduced from the pressure of the reactor of the second stage to the pressure of the reactor of the third stage by means of a regulating valve or some other device characterized by a pressure drop.

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- 11. A process according to claim 1, wherein the 25 reactor of the first stage is integrated into the reactor of the second stage.
- 12. A process according to any of claims 1 to 10, wherein the pressure upstream of the static mixer is 3-30 70 bar, preferably 15-45 bar, the pressure in the reactor of the second stage is 2.5-35 bar, preferably 15-35 bar, and the pressure in the reactor of the third stage is 2-20 bar, preferably 3.5-16 bar.
- 35 13. A process according to any of claims 1 to 11, wherein the temperature in the first, second and third stages is in each case 80-190°C, preferably 90-150°C.

- 14. A process according to any of claims 1 to 12, wherein an aromatic hydrocarbon such as toluene or preferably a chlorinated aromatic hydrocarbon such as chlorobenzene, ortho-dichlorobenzene or trichlorobenzene or a mixture thereof is used as inert solvent.
- 10 No pages of drawings follow

Continuous preparation of isocyanates

Abstract

The invention relates to a process for preparing polyisocyanates by reacting organic amines with phosgene, wherein the reaction is carried out in at least three stages, with the first stage being carried out in a mixing apparatus, the second stage in at least one residence apparatus and the third stage in at least one separation apparatus and the pressure in each successive stage being lower than that in the previous stage.